SepraPor[®] 50 kDa Hollow Fiber Tangential Flow Filter Cartridge

Description

SepraPor® XFM050 is a tangential flow filtration (TFF) cartridge containing hollow fiber membranes. It is ideal for use in a variety of biopharmaceutical applications including ultrafiltration, diafiltration, and purification. This TFF cartridge is available in a range of sizes from bench through production scale for ease of scalability. The SepraPor® filter cartridge is 100% integrity tested during manufacture and meets the critical demands of the pharmaceutical, biotechnology, and related industries.

Materials of Construction

All components of the filter are animal component free (ACF). These materials are listed for food contact use in the Code of Federal Regulations (CFR), Title 21, as below:

Hollow fibers:	Polysulfone CFR Title 21, 177.1		
Fiber bundle netting:	Polypropylene CFR Title 21, 177.1		
Outer sleeve:	Polysulfone	CFR Title 21, 177.1655	
Fiber encapsulation:	Ероху	CFR Title 21, 175.300	
O-rings:	Silicone	CFR Title 21, 177.2600	
Retention Rating	50 kDa		
Typical Fiber Lumen	1.0 mm		
Fluid Path Length (Nominal)	12 in (30 cm)		
2 . ,	24 in (60 cm)		
Maximum Diffusive Flow Rate	2.79 mL/min/m ² @ 30 psig (2.07 bar), water with air		
Minimum Permeate Flux	100 - 480 LMH/bar (Liters/m²/h/bar) @ 25 °C		
Operating Characteristics			
Operating temperature range:	32 °F to 100 °F (0 °C to 38 °C)		
Maximum operating temperature:	122 °F (50 °C) for short-term operations such as cleaning		
Maximum feed pressure:	65 psig @ 77 °F (4.48 bar @ 25 °C)		

Sterilization

Autoclave or steam in place (SIP): 121 °C to 123 °C (15 psi, 1.03 bar), 30 minutes. Apply a validated sterilization cycle that increases and decreases temperature gradually. Consult sterilization instructions in the SepraPor[®] Green Docs document for further guidance.

Maximum transmembrane pressure: 45 psig @ 77 °F (3.10 bar @ 25 °C)

Biological Safety

SepraPor[®] filters meet the requirements as specified in the USP 43 Biological Reactivity Tests, *in vitro* <87> (cytotoxicity), and polysulfone and epoxy meet *in vivo* <88> (Class VI Plastics).

Quality Assurance

SepraPor[®] filters comply with the Food and Drug Administration Code of Federal Regulations, Title 21, Parts 210 and 211. Product is manufactured and packaged in a cleanroom facility that, through voluntary compliance, meets or exceeds FDA Good Manufacturing Practice Standards. To ensure product reliability, Meissner's Quality Assurance staff continually audits the manufacturing process for conformance to its Quality Management System. Each SepraPor[®] filter is labeled with filter type, lot number, and unique serial number. The serial number for all cartridge filters can be found on the product packaging.



Ordering Guide

Part Number	Filter Diameter	Fluid Path Length (Nominal)	Effective Filtration Area	Housing Seal Configuration	O-ring Seal Material
XFM050C412-AAS	2" (5.1 cm)	12" (30 cm)	4.3 ft ² (0.4 m ²)	AA	Silicone
XFM050C424-AAS		24" (60 cm)	8.6 ft ² (0.8 m ²)		
XFM050C512-AAS	3" (7.6 cm)	12" (30 cm)	9.7 ft ² (0.9 m ²)		
XFM050C524-AAS		24" (60 cm)	22.6 ft ² (2.1 m ²)		

Additional information about this filter product is available in the SepraPor® Green Docs document at www.meissner.com/green-docs.

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